

Finite Element Solution of the Steady-State Smoluchowski Equation for Rate Constant Calculations

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ABSTRACT This article describes the development and implementation of algorithms to study diffusion in biomolecular systems using continuum mechanics equations. Specifically, finite element methods have been developed to solve the steady-state Smoluchowski equation to calculate ligand binding rate constants for large biomolecules. The resulting software has been validated and applied to mouse acetylcholinesterase. Rates for inhibitor binding to mAChE were calculated at various ionic strengths with several different reaction criteria. The calculated rates were compared with experimental data and show very good agreement when the correct reaction criterion is used. Additionally, these finite element methods require significantly less computational resources than existing particle-based Brownian dynamics methods.

INTRODUCTION

Diffusion plays a central role in numerous biological processes, governing the kinetic properties of events across a variety of length scales: from ligand binding (Antosiewicz et al., 1995, 1996b; Antosiewicz and McCammon, 1995; Lesyng and McCammon, 1993; McCammon and Karplus, 1977; Northrup et al., 1984; Tan et al., 1993; Tara et al., 1998; Wade et al., 1994) to protein-protein encounter (Elcock et al., 2001; Gabdoulline and Wade, 2001; Northrup and Erickson, 1992; Sheinerman et al., 2000; Zhou, 1997) to signal transmission at synaptic junctions (Franks et al., 2002; Kara and Friedlander, 1998; Roberts, 1994; Smart and McCammon, 1998; Tai et al., 2003; Zoli and Agnati, 1996). Biological simulations have been used to study such diffusion-controlled processes in a number of settings and have provided useful insight into the molecular determinants of the kinetic parameters. However, accurate modeling of diffusion within biomolecular systems while incorporating the effects of ionic strength, solvent, and protein charges, and applying to large biological systems with complex geometries, has proven to be the rate-limiting step for a variety of such simulations.

Currently, standard techniques for modeling diffusional processes can be loosely grouped into *particle-based* and *continuum* methods. Particle-based methods are typically stochastic in nature and include Monte Carlo (Berry, 2002;

Genest, 1989; Saxton, 1992; Stiles and Bartol, 2000; Brownian dynamics (BD) (McCammon, 1987; Northrup et al., 1988a; Wade et al., 1993), and Langevin dynamics (Eastman and Doniach, 1998; Yeomans-Reyna and Medina-Noyola, 2001) simulations. The connection between BD simulations and of the calculation of association rate constants was established by Northrup, Allison, and McCammon (Northrup et al., 1984) and has been studied by numerous others (Antosiewicz et al., 1996a; Antosiewicz and McCammon, 1995; Chung et al., 2002; Northrup et al., 1988b; Tan et al., 1993; Tara et al., 1998; Wade et al., 1993; Zhou, 1993; Zhou et al., 1998a; Zhou and Szabo, 1996; Zou et al., 2000). In contrast to particle-based approaches, continuum methods describe diffusional processes in terms of probability or concentration profiles rather than simulating the stochastic motion of individual particles. Continuum methods are typically based on solutions of partial differential equations such as the diffusion or Smoluchowski equation (Chan and Halle, 1984; Gardiner, 1997; Lenzi et al., 2003; Smart and McCammon, 1998; Tai et al., 2003); these solutions can then be processed to determine ligand-protein binding (Agmon et al., 1991; Smart and McCammon, 1998; Tai et al., 2003; Zhou, 1990) or dissociation (Agmon, 1984). These methods have been particularly popular in the fields of ion channel (Coalson and Duncan, 1992; Gillespe et al., 2002; Im and Roux, 2002; Kurnikova et al., 1999a) and semiconductor (Selberherr, 1984) modeling.

Both particle-based and continuum diffusion methods have their relative strengths. Particle-based methods can deal with a wide range of diffusing molecular geometries, whereas continuum methods are restricted to spherical ligands. This spherical approximation is likely to be most appropriate for substrates with charge distributions with small multipole moments and reaction criteria that do not require

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a detailed fit between the substrate and macromolecule. Additionally, particle-based approaches permit the natural inclusion of stochastic reaction phenomena and other complicated boundary conditions. However, the stochastic nature of particle-based approaches can lead to convergence problems which are not present in the deterministic continuum method. Furthermore, continuum approaches facilitate the inclusion of other continuum phenomena such as elastic deformations and fluid flow. Finally, as illustrated in this article, the computational cost of the continuum simulations is significantly smaller than for particle-based methods.

Currently, there are a number of tools available for particle-based biomolecular diffusion simulations, including: SDA (Gabdouline and Wade, 1997), UHBD (Briggs et al., 1995), MacroDox (Northrup et al., 1999), and MCell (Stiles and Bartol, 2000). However, there are no biomolecule-specific tools available for analyzing diffusion via continuum models and only a few general diffusion tools (Krissinel and Agmon, 1996). The objective of this study is to develop, validate, and apply algorithms to solve the steady-state Smoluchowski equation (SSSE) with finite element methods using realistic biomolecular geometries to determine the steady-state ligand binding rate constant. Specific aims in this study include: development of the adaptive meshing method to realistically describe biomolecular geometries; development of the finite element solver of the steady-state Smoluchowski equation to analyze the concentration of the diffusing particles and calculate the association rate constants; validation of the SSSE with a simple spherical biomolecular system through the comparison with the analytical results; and application of the validated SSSE solver to mouse acetylcholinesterase (mAChE) ligand binding.

THEORY AND ALGORITHMS

The steady-state Smoluchowski equation

The Smoluchowski equation describes the overdamped (i.e., instantaneous momentum relaxation) dynamics of multiple particles while neglecting interparticle interactions (Smoluchowski, 1917; Szabo et al., 1988; Zhou, 1990). For a stationary diffusion process, the Smoluchowski equation has the steady-state form of

$$Lp(x) = \nabla \cdot D(x)[\nabla p(x) + \beta p(x)\nabla W(x)] = 0, \quad (1)$$

where $Lp(x)$ represents $(dp(x, t)/dt)$ (t is the time), $p(x)$ is the distribution function of the reactants, $D(x)$ is the diffusion coefficient, $\beta = 1/kT$ is the inverse Boltzmann energy, k is the Boltzmann constant, T is the temperature, and $W(x)$ is the potential mean force (PMF) for the diffusing particle. The above steady-state Smoluchowski equation (SSSE) can also be written in terms of the flux operator J , which generates vector-valued functions and is defined as

$$Jp(x) = D(x)[\nabla p(x) + \beta p(x)\nabla W(x)], \quad (2)$$

allowing Eq. 1 to be rewritten as

$$Lp(x) = \nabla \cdot Jp(x) = 0. \quad (3)$$

The SSSE can be solved to determine bimolecular diffusional encounter rates. Following the work of Zhou (1990), the application of the SSSE to this problem involves the solution of Eq. 3 in a three-dimensional domain Ω , with the following boundary conditions: a bulk Dirichlet condition on the outer boundary $\Gamma_b \subset \partial\Omega$,

$$p(x) = p_{\text{bulk}} \quad \text{for } x \in \Gamma_b, \quad (4)$$

specifying the bulk concentration p_{bulk} ; a reactive Robin or Dirichlet condition on the active site boundary $\Gamma_a \subset \partial\Omega$,

$$n(x) \cdot Jp(x) = \alpha(x)p(x) \quad \text{for } x \in \Gamma_a, \quad (5)$$

or

$$p(x) = 0 \quad \text{for } x \in \Gamma_a, \quad (6)$$

providing either an intrinsic reaction rate $\alpha(x)$ or an absolute reactivity, respectively; and a reflective Neumann condition on the nonreactive boundary $\Gamma_r \subset \partial\Omega$,

$$n(x) \cdot Jp(x) = 0 \quad \text{for } x \in \Gamma_r. \quad (7)$$

The problem domain is depicted in Fig. 1; Δ is a simply connected domain with boundary Γ_b , which represents the volume containing the reactive object and the solvent. The domain $\Xi \subset \Delta$ is a simply connected region representing the reactive object with boundary $\Gamma_{\text{ar}} = \Gamma_a \cup \Gamma_r$ such that $\Gamma_a \cup \Gamma_r = \mathbf{0}$. The Γ_r portion of this boundary represents the

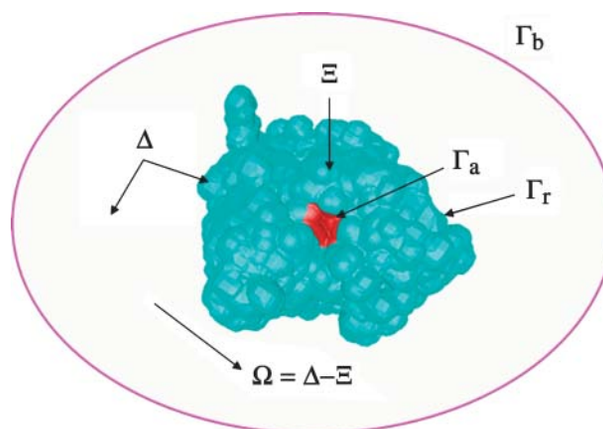


FIGURE 1 Schematic of problem domain denoting the various surfaces and volumes described in the text.

nonreactive surface of the biomolecular object whereas the Γ_a boundary represents the reactive region of the object. The problem domain is the region in which substrate is allowed to diffuse, namely $\Omega = \Delta - \Xi$. To ensure the well-posedness of the SSSE Smoluchowski equation, we place several physically based restrictions on the problem data: the potential of mean force (PMF, the effect interaction potential) is bounded and dies off with increasing distance, the diffusion constant is positive definite and finite, and the temperature is positive definite and finite. Finally, the diffusion-influenced biomolecular reaction rate constant is obtained from the flux by integration over the active site boundary, as

$$k = \frac{\int_{\Gamma_a} n(x) \cdot Jp(x) ds}{p_{\text{bulk}}}. \quad (8)$$

Adaptive mesh generation

One of the challenges in solving the SSSE is the development of meshes which respect the complicated biomolecular geometry. Not surprisingly, the results of finite element solution of the SSSE are sensitive to the quality of the finite element discretization, therefore robust methods must be used to generate the adaptive meshes. The geometry of the mesh is set by the underlying arrangement of atoms within the stationary biomolecule. This atomic geometry is transformed into a scalar accessibility function in a manner analogous to that used for Poisson-Boltzmann electrostatics calculations (Baker et al., 2000, 2001). Specifically, we use a characteristic function $\chi(x)$ which represents an inflated van der Waals-based accessibility, as

$$\chi(x) = \begin{cases} 0, & \text{if } \|x - y_i\| < r_i + \sigma \text{ for } i = 1, \dots, N, \\ 1, & \text{otherwise} \end{cases}, \quad (9)$$

where (y_i, r_i) are the coordinates and radii of the N atoms in the biomolecule and σ is the radius of the diffusing species. This accessibility function provides an abstraction of the biomolecular structure which can then be used as input for advanced volume isocontouring methods. Specifically, $\chi(x)$ provides a (grid-based) dataset which is then isocontoured (at a value of 0.5) via dual contouring methods into a three-dimensional tetrahedral mesh (Zhang et al., 2003b).

The goal of the dual contouring methods used in this work is to tetrahedralize the interval volume between the biomolecular surface and an outer boundary sphere S_1 which is usually 40 times that of the biomolecule (compare to Figs. 2 and 3). The resulting tetrahedral mesh is spatially adaptive and preserves molecular surface features while minimizing the number of simplices. The four main steps of our adaptive tetrahedral meshing of the problem domain are described in the following sections.

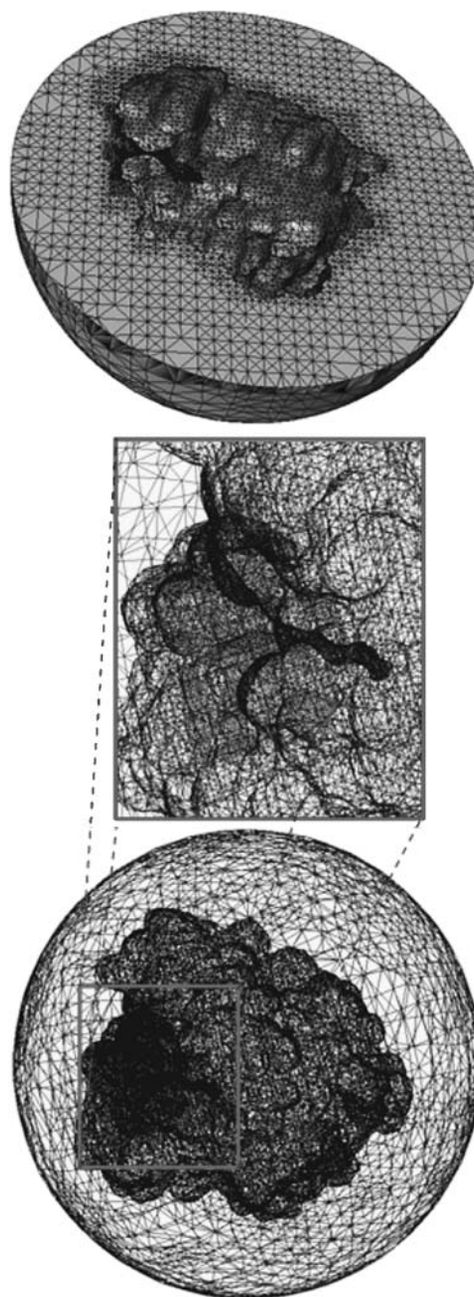


FIGURE 2 Adaptive tetrahedral meshes for mouse acetylcholinesterase. (Bottom) The molecular surface and outer sphere S_0 ; the active site gorge is shown in greater detail inside the red box. (Middle) Magnification of the red box in the bottom picture. (Top) The tetrahedral mesh of the interval volume between the molecular surface and the outer sphere S_0 (cross section).

Data rescaling

We select a sphere S_0 with a radius (r_0) which is larger than the biomolecular radius, and add it outside the biomolecular surface. For each data point inside the molecular surface, we keep the original function value whereas for each data point outside the molecular surface, we reset the function value as the smaller of the original function value and the shortest

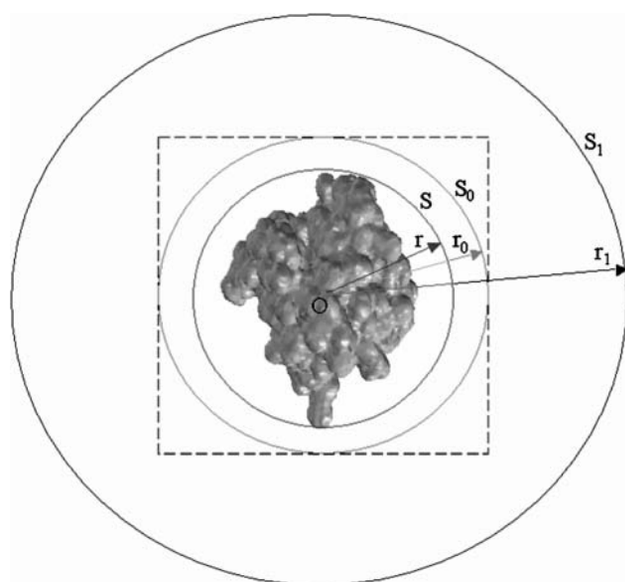


FIGURE 3 Data scaling for the adaptive mesh; surfaces are described in the text.

distance from the grid point to the sphere S_0 . We tetrahedralize the interval volume between the biomolecular surface and an outer boundary sphere S_0 , and then extend the tetrahedral mesh to the sphere S_1 , shown in Fig. 3.

Tetrahedral mesh extraction

Dual contouring (Ju et al., 2003) uses an octree data structure to analyze mesh edges, called *sign-change edges*, which have endpoints lying on different sides of the isosurface of interest. The mesh adaptivity is determined during a top-down octree construction. Each sign change edge is shared by either three (adaptive case) or four (uniform case) octree cells. A minimizer point is calculated for each cell by minimizing a predefined quadratic error metric (Garland and Heckbert, 1998). Finally, for each sign change edge, a quadrilateral or triangle is constructed by connecting the minimizer points; these resulting quadrilaterals and triangles provide a dual piecewise linear approximation of the isosurface. This dual contouring method has already been extended to extract tetrahedral meshes from volumetric scalar fields (Zhang et al., 2003b) following a similar procedure to the two-dimensional case.

Three-dimensional dual contouring is used to construct the tetrahedral mesh between the biomolecular surface and the sphere S_0 . When used as described above, the resulting mesh is finest around the molecular surface, and gradually gets coarser away from the molecule, toward the boundary S_0 . However, the adaptive nature of this method can also be exploited to provide additional detail in biologically important regions of the molecule such as active sites. As illustrated in Figs. 2 and 6 for mouse acetylcholinesterase, we control the mesh resolution to generate a very fine mesh

in the active site gorge region and a relatively coarse mesh elsewhere near the biomolecular surface.

Extension to the outer boundary

This initial tetrahedral mesh is modified by gradually adding tetrahedra of increasing size from the boundary of sphere S_0 to the outer boundary to produce a finite element mesh spanning the entire problem domain. Care is taken when adding these exterior tetrahedra to ensure the quality of the simplex shape and mesh topology.

Quality detection and improvement

The tetrahedral mesh generated by the dual contouring method may have isolated vertices, nonsimple components, or overlapping tetrahedra—problems which must be removed before the mesh can be used to solve the SSSE. Isolated vertices and nonsimple components are identified and removed using the connectivity information (i.e., edges and simplices) intrinsic to the finite element mesh. Overlapping simplices are corrected using methods previously described by Zhang et al. (2003b).

In addition to correcting these topological problems, the geometric quality of the mesh must be checked and, if necessary, improved. Poor quality tetrahedra called “slivers” are common in most tetrahedral mesh generation methods. The presence of such slivers can often confound finite element solvers and therefore these poor-quality tetrahedra must be removed (Cheng and Dey, 2002; Cheng et al., 2000) before using the generated meshes to solve the SSSE. We combine edge contraction and smoothing methods to improve the quality of the meshes based on tetrahedral-edge ratios, Joe-Liu shape metrics (Liu and Joe, 1994), and minimum volume bounds. Edge contraction (Cheng and Dey, 2002; Cheng et al., 2000) improves tetrahedral shape by merging vertices to combine smaller simplices into larger, better quality tetrahedra. Additionally, we use a smoothing method based on multilinear averaging (Zhang et al., 2003b) to improve the Joe-Liu shape metric and the minimum volume bounds.

Finite element discretization of the Smoluchowski equation

To solve the SSSE numerically as a finite system, it is necessary to truncate and discretize the infinitely large problem domain implicit in Eqs. 1–7. We solve Eq. 1 using finite element methods (Axelsson and Barker, 2001; Braess, 1997) inside a domain Ω that is 40 times the scale of the size of the biomolecule. Since the effects of the PMF are not included beyond the outer boundary, this large size is typically necessary for electrostatic forces to decay to ~ 0 and/or avoid more complicated outer boundary conditions. However, due to the adaptive nature of the finite

element meshes used to discretize the system and our multiresolution approach to the calculation of the potential mean force (see below and Baker et al., 2001), this large mesh does not impose a significant computational burden. The outer boundary of the diffusion domain Ω is subject to the Dirichlet boundary condition per Eq. 4. The inner (molecular) boundary is assigned reactive and reflective conditions as shown in Fig. 1 and described in Eqs. 5–7. The whole domain is discretized on an adaptively generated tetrahedral mesh using the methods outlined above.

The resulting tetrahedral mesh forms the structure over which we define a function space $V_h = \text{span}\{v_i\}$, where $\{v_i\}$ is the set of piecewise-linear finite element basis functions defined over each tetrahedral vertex. The solution to the SSSE is approximated by a function, $p_h \in \bar{p}_h + V_h$, constructed from the linear combination of basis functions,

$$p_h(x) = \sum_i a_i v_i(x). \quad (10)$$

The trace function \bar{p}_h is not explicitly constructed, but is assumed to satisfy the Dirichlet boundary conditions in Eqs. 4 and 6. For this construction of p_h from piecewise-linear functions to be successful, we must restate the SSSE equations in their weak form. Clearly, piecewise-linear functions do not provide a well-defined second derivative, as required by Eq. 1. This difficulty can be overcome by integrating the SSSE with a test function $v(x)$, as

$$\int_{\Omega} v(x) \nabla \cdot Jp(x) dx = 0. \quad (11)$$

Integrating the above equation by parts via Green's theorem gives

$$\int_{\Omega} \nabla v(x) \cdot Jp(x) dx - \int_{\partial\Omega} v(s) Jp(s) \cdot n(s) ds = 0, \quad (12)$$

and incorporation of the various boundaries conditions Eqs. 4–7, allows the boundary integral to be simplified as

$$\int_{\Omega} \nabla v(x) \cdot Jp(x) dx - \int_{\Gamma_a} v(s) \alpha(s) p(s) ds - \int_{\Gamma_b} v(s) J\bar{p}(s) \cdot n(s) ds = 0 \quad (13a)$$

or

$$\int_{\Omega} \nabla v(x) \cdot Jp(x) dx - \int_{\Gamma_a \cup \Gamma_b} v(s) J\bar{p}(s) \cdot n(s) ds = 0, \quad (13b)$$

depending on the choice of reactive boundary condition (compare to Eq. 5 and Eq. 6, respectively). Finally, the above

integrals can be rewritten in terms of the bilinear form $\langle F(p), v \rangle$,

$$\langle F(p), v \rangle = \int_{\Omega} \nabla v(x) \cdot Jp(x) dx - \int_{\Gamma_a} v(s) \alpha(s) p(s) ds - \int_{\Gamma_b} v(s) J\bar{p}(s) \cdot n(s) ds \quad (14a)$$

$$\langle F(p), v \rangle = \int_{\Omega} \nabla v(x) \cdot Jp(x) dx - \int_{\Gamma_a \cup \Gamma_b} v(s) J\bar{p}(s) \cdot n(s) ds, \quad (14b)$$

to give the weak form of the SSSE,

$$\text{Find } p_h \in \bar{p}_h + V_h \text{ such that } \langle F(p_h), v_i \rangle = 0 \text{ for all } v_i \in V_h. \quad (15)$$

This form of the SSSE requires only one order of differentiation under an integral and is therefore a weaker formulation of the SMOL equation than the original second-order differential Eq. 1. Equation 15 is the foundation for the finite element solvers used in this work.

In the Smoluchowski equation, the diffusion coefficient $D(x)$ and electrostatic potential mean force $W(x)$ all are spatially dependent. Ignoring hydrodynamic interactions (which could be included as part of the spatial dependence of D), the diffusion coefficient is treated as a constant. In most BD models, the electrostatic potential is treated independently from the diffusing species—eliminating the possibility of screening by ligand, substrate inhibition, etc. This is also the model we will employ for these initial studies, implying that Eq. 15 is a linear equation. Discretization of Eq. 15 with a finite element basis (per Eq. 10) leads to a linear system of (sparse) equations which can be solved using standard linear algebra methods. However, it is important to note that this model could easily be extended to the related Poisson-Nernst-Planck (PNP) system where the PMF is coupled to the diffusing species. For PNP equations, well-established nonlinear methods will need to be employed (Holst and Saied, 1995; Koumanov et al., 2003; Kurnikova et al., 1999b; Schuss et al., 2001). Additionally, like many PNP methods, the present model does not consider correlations between diffusing species. Such effects have been shown to be important in confined spaces such as ion channels (Boda et al., 2002; Gillespe et al., 2002) or at higher concentrations of multivalent ions (Holm et al., 2001). For the low concentrations of ligand used in many protein-ligand diffusive encounter simulations, these correlation effects are likely to be relatively small; however, it is important that such effects are considered when performing diffusion simulations. Finally, the present model does not include dielectric boundary or apolar forces which have been shown

to be important factors for diffusion behavior in some biological systems (Nadler et al., 2003). Such effects are very important for diffusion of charged species in confined spaces (e.g., ion channels) but have a much less significant impact for ligand-protein binding at relatively low concentrations (where dielectric boundary and apolar forces often cancel).

Implementation

We have developed a software package called SMOL which uses the finite element method to solve the SSSE for biomolecular systems. The SMOL program is based on the LBIE-Mesh software (<http://www.ices.utexas.edu/CCV/software>), FETk finite element software library (M. Holst, Adaptive Multilevel Finite Element Methods on Manifolds and their Implementation in FETk; in preparation; currently available as a technical report and User's guide to the FETk software; see <http://www.fetk.org/>) and the APBS software package (<http://agave.wustl.edu/apbs/>, Baker et al., 2001). Specifically, SMOL uses LBIE-mesh for construction of the initial mesh, FETk's finite element infrastructure for the discretization and solution of the linear system of equations. APBS is used to calculate the electrostatic component of the potential mean force by solution of the Poisson-Boltzmann equation. The SMOL software is still actively under development and will be released under the open-source GNU public license (see <http://www.gnu.org/>). Adventurous readers are welcome to contact the authors for access to early versions of the software; others should visit <http://agave.wustl.edu/> for more information about user-friendly public releases.

The sequence of operations followed by the SMOL program during a typical solution of the SSSE for biomolecular rate calculation is outlined in the following sections.

Parameterization

The biomolecular coordinates from the Protein Data Bank must be parameterized with appropriate atomic radii and charges. The current work uses the CHARMM22 force field (Brooks et al., 1983); conversion tools for several other force fields are provided with the APBS software or available through the PDB2PQR web service (<http://nbcv.sdsc.edu/pdb2pqr/>).

Mesh construction

First, a grid-based form characteristic accessibility function $\chi(x)$ is generated with APBS using the diffusing species' radius σ (compare to Eq. 9) and the parameterized biomolecular structure as input data. This discretized characteristic function is then used with the adaptive meshing tools described above to generate quality tetrahedral meshes of the problem domain. Finally, boundary conditions as in Eqs. 4–7 are assigned to the mesh based on user-defined spatial

criteria using knowledge of the active site location and other structural features.

Potential of mean force

Only electrostatic contributions to the PMF are included in this version of the software. The electrostatic potential is calculated over the entire diffusion domain of the biomolecule with APBS. This data is stored in a parallel, multiresolution format where (sometimes multiple) high-resolution meshes provide PMF values near the molecular surface while coarser meshes provide values away from the biomolecule. This decomposition of the potential closely follows the parallel-focusing methods described elsewhere (Baker et al., 2001).

Solution of the SSSE and analysis

Given the above PMF data, finite element mesh, and user-defined values for the diffusing particle's charge, concentration, and diffusion constant data, the SSSE is solved with the finite element methods described above. After the successful solution of the differential equation, output is provided for the steady-state rate constant and (if requested) the concentration profile of ligand around the biomolecule.

VALIDATION OF THE SMOL PROGRAM WITH A SPHERICAL TEST CASE

Before applying the SMOL program to a biomolecular system with complex geometry, we first tested it with the classic spherical system (Krissinel and Agmon, 1996) and compared the calculated result with the known analytical solution. For this test case, we chose a fixed sphere with an 8 Å radius and +1 e charge and a diffusing sphere with a 2 Å radius and variable charge. The partial domain for this spherical molecule is shown in Fig. 4 A. The entire problem domain was discretized with 1,024,752 tetrahedral elements. A detailed view of the surface mesh for the stationary sphere is also shown in Fig. 4 B. While solving the Smoluchowski equation, the whole surface of biomolecule sphere Γ_a was treated with a perfectly absorbing zero Dirichlet reactive boundary condition as Eq. 6. The diffusing particle's dimensionless bulk concentration was set to 1. Ignoring hydrodynamic interactions, the diffusion constant D is calculated as $7.8 \times 10^4 \text{ \AA}^2/\mu\text{s}$ using the Stokes-Einstein equation with a hydrodynamic radius of 3.5 Å, solvent viscosity of $0.891 \times 10^{-3} \text{ kg/(m s)}$, and 298 K temperature.

Analytical solution

For a spherically symmetric system with a Coulombic form of the PMF, $W(r) = q/r$, the SSSE can be written as

$$\frac{1}{r^2} \frac{\partial}{\partial r} (r^2 J_p) = \frac{1}{r^2} \frac{\partial}{\partial r} \left(r^2 D \left(\frac{\partial p}{\partial r} - \beta p \frac{q}{r^2} \right) \right) = 0, \quad (16)$$

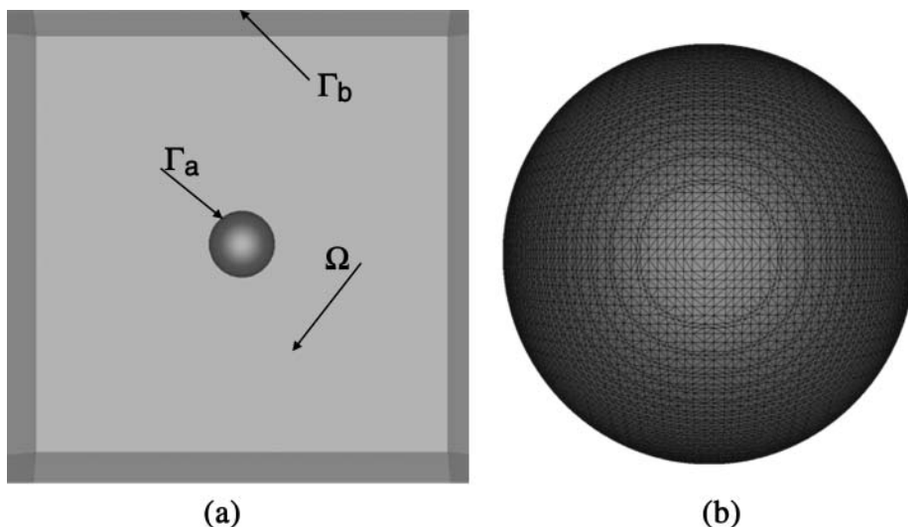


FIGURE 4 Illustration of the discretized problem domain for the spherical test case. (a) Partial domains of the fixed sphere; the outer boundary of the domain is 40 times the radius of the sphere. (b) Subset of the mesh near the fixed sphere surface.

with boundary conditions

$$p(r_1) = 0, p(r_2) = p_{\text{bulk}}, \quad (17)$$

where r_1 and r_2 are the radii for the reactive and outer boundaries. The analytical expression for the association rate constant obtained from the solution of Eqs. 16 and 17 is

$$k = 4\pi r_1^2 Jp(r_1) = 4\pi Dqp_{\text{bulk}} \left(\frac{e^{q/r_1}}{e^{q/r_1} - e^{q/r_2}} \right). \quad (18)$$

This analytical form of the solution was evaluated using the parameters listed above with varying ligand charges in the infinite dilution limit: $r_2 \rightarrow \infty$. Table 1 presents the numerical values of these analytical reaction rates as a function of effective ligand charge.

SMOL numerical solution

Reaction rates were also calculated with the SMOL software using the parameters given above and a finite problem domain with the outer boundary at 400 \AA ; this domain was discretized with 1,024,752 tetrahedral elements. The results of these calculations are shown in Table 1. The calculated reactive area is 1257.98 \AA^2 , differing by only 0.11% from the analytical area of 1256.64 \AA^2 and illustrating that the finite element mesh realistically represents the system's geometry.

The performance of the SMOL program is good, with a typical relative error of 1% and the largest error (2.7%) found for the more challenging mutually repulsive cases.

APPLICATION OF THE SMOL PROGRAM TO MOUSE ACETYLCHOLINESTERASE

One of the major advantages of continuum methods such as the SSSE is the ability to simulate diffusion to large biological systems with complex geometries with significantly lower computational cost than Brownian dynamics techniques. This section demonstrates the use of SSSE to study the ligand binding kinetics of acetylcholinesterase (AChE, E.C. 3.1.1.7) (Quinn et al., 1995). AChE is a serine esterase that terminates the activity of acetylcholine (ACh) within the cholinergic synapse by hydrolysis of the ACh ester bond to produce acetate and choline (Berg et al., 1995). Hydrolysis of ACh occurs in the active site of AChE, which lies at the base of a 20 \AA -deep gorge within the enzyme. The rate-limiting step of ACh hydrolysis by AChE is the diffusional encounter (Bazelyansky et al., 1986; Berman et al., 1991; Nolte et al., 1980), making the system a popular target for both experimental (Bourne et al., 1995; Radic et al., 1997; Velsor et al., 2003) and computational diffusion studies (Tan et al., 1993; Tara et al., 1998).

To provide additional data for assessment of SMOL and the SSSE in rate constant calculations, we based our analysis of AChE kinetics on previous BD work by Tara et al. (1998). Specifically, we used a mouse AChE (mAChE) structure

TABLE 1 Analytical and numerical reaction rates for the spherical system as a function of effective ligand charge

Ligand charge (e)	1.0	0.8	0.6	0.4	0.2	0.0	-0.2	-0.4	-0.6	-0.8	-1.0
Analytical results $k_{\text{on}} (10^{11} \text{ M}^{-1} \text{ min}^{-1})$	2.41	2.61	2.82	3.05	3.28	3.62	3.79	4.06	4.34	4.64	4.95
SMOL results $k_{\text{on}} (10^{11} \text{ M}^{-1} \text{ min}^{-1})$	2.44	2.63	2.83	3.04	3.26	3.49	3.74	3.99	4.25	4.53	4.81

adopted from the crystal structure of the mAChE-fasciculin 2 complex (1MAH) (Bourne et al., 1995) and perturbed by Tara and co-workers via molecular dynamics simulations with an ACh-like ligand in the active site gorge (Tara et al., 1998) to produce gorge conformations with wider widths than the original x-ray structure. The diffusing ligand was modeled as a sphere with an exclusion radius of 2.0 Å and a diffusion constant of $7.8 \times 10^4 \text{ Å}^2/\mu\text{s}$. This perturbation was necessary for computational diffusion simulations with a fixed biomolecular structure. Related work (Baker and McCammon, 1999; Zhou et al., 1998b) supports the use of this open gorge state due to its relevance in conformational gating dynamics of the enzyme. The geometry of mAChE and its active site gorge are shown in Figs. 5 and 6.

Reactive boundary definitions

In our calculation of binding rates with SMOL, reactive boundaries were defined following the typical BD methods: a spherical reactive surface is defined at an arbitrary radius from the biomolecular active site (see Fig. 6). To test the sensitivity of the results to the positions of the reactive boundary, six reactive surfaces were placed along the gorge and the gorge opening. Following Tara et al. (1998), these surfaces were based on six spheres placed along the active site gorge (in what follows, the coordinates are defined with the carbonyl carbon of residue S203 in the mAChE structure at the origin and the gorge aligned with the y axis): sphere 1 centered at (0.0, 16.6, 0.0) with a 12 Å radius; sphere 2 centered at (0.0, 13.6, 0.0) with a 9 Å radius; sphere 3 centered at (0.0, 10.6, 0.0) with a 6 Å radius; sphere 4 centered at (0.0, 7.6, 0.0) with a 6 Å radius; sphere 5 centered at (0.0, 4.6, 0.0) with a 6 Å radius; and sphere 6 centered at (0.0, 1.6, 0.0) with a 6 Å radius. Each reactive surface N is defined by explicitly including the union of spheres N through 6 in the mAChE structure—the actual reactive surface is then simply that portion of the new “molecular” surface due to the intersection of added spheres with molecule. The six reactive surfaces based on this spherical definition are shown in Fig. 6.

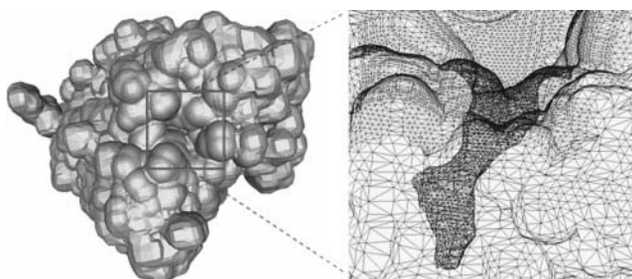


FIGURE 5 Geometry of mAChE finite element mesh (*left*) and its active site gorge (*right*).

Variables and parameters for the biomolecule and diffusion domain

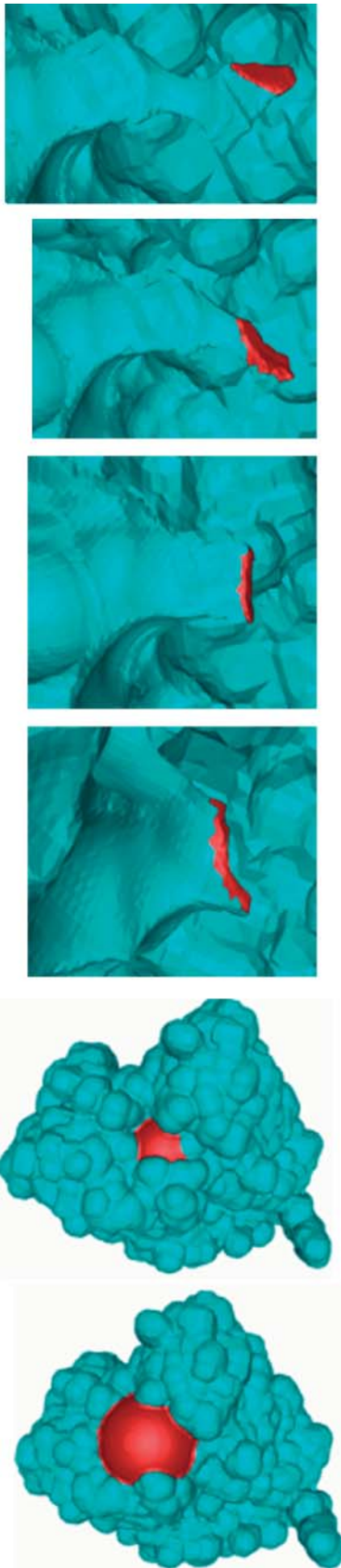
For the purpose of the PMF (electrostatics) calculation for mAChE, partial charges and radii were assigned from the CHARMM22 force field. Poisson-Boltzmann electrostatics calculations were performed using dielectric values of 4 and 78 for the protein and solvent, respectively; a solvent probe radius of 1.4 Å; and an ion exclusion layer of 2.0 Å. APBS was used to solve the nonlinear Poisson-Boltzmann equation using the parallel focusing method (Baker et al., 2001). Sets of nested potential grids were obtained with the finest grid having dimensions $76 \times 66 \times 91 \text{ Å}$ with 161 grid points in each direction. For the study of the effect of ionic strength on reaction rate, separate calculations were performed at ionic strengths of 0.00, 0.050, 0.100, 0.150, 0.200, 0.250, 0.300, 0.450, 0.600, and 0.670 M.

The problem domain definition for the mAChE system is shown in Fig. 1. The outer boundary Γ_b has a radius of 40 times the size of mAChE molecule, and is given a bulk concentration of unity. Reactive surface Γ_a is assigned zero (perfectly absorbing) Dirichlet boundary conditions (compare to Eq. 6) and the remainder of the biomolecular surface is treated as the reflective boundary Γ_r with Neumann condition (Eq. 7). The adaptive meshing method described above was used to discretize the problem domain. To accurately reproduce the topology of the 20 Å-deep active site gorge, a finer mesh was used for discretization near catalytic site and gorge; other areas of the molecule are discretized with relatively coarser mesh. The resulting mesh contained 656,823 simplices and 121,670 vertices. A subset of the domain is shown in Figs. 3, 5, and 6 to illustrate the general features of the mesh. Detailed views of the mesh near the active site gorge are shown in Figs. 5 and 6.

The diffusing particle was treated as a sphere with $a + 1 e$ charge, a 2.0 Å exclusion radius, and a diffusion constant of $7.8 \times 10^4 \text{ Å}^2/\mu\text{s}$ (parameters were taken from Tara et al., 1998, which are similar to those for the TFK⁺ ligand).

Calculating the reaction rate for mAChE-TFK⁺ encounter with SMOL

Reaction rates for mAChE-TFK⁺ diffusional encounter were calculated from the SSSE via the SMOL program for ionic strengths ranging from 0.000 to 0.670 M NaCl and for the reactive boundary definitions described above. All calculations were performed on Intel Pentium 4 CPU 2.20-GHz computer systems with 1.5 GB RAM running RedHat Linux 9. SMOL executables were compiled with the Intel FORTRAN and C compilers using the $-O2$ optimization. The SMOL calculations took an average of 55 s per reactive surface. The resulting reaction rates are presented in Fig. 7 and Table 2; analysis of this data and comparison with both experimental and BD results are presented below.



Calculating the reaction rate for mAChE-TFK⁺ encounter with UHBD

The BD simulations of Tara et al. (1998) were repeated using the UHBD software (Briggs et al., 1995) to obtain timing data and raw reaction rates for comparison with the SMOL results. As for the SMOL calculations, all simulations were performed on Intel Pentium 4 CPU 2.20-GHz computer systems with 1.5 GB RAM running RedHat Linux 9; the UHBD executables were compiled with the Intel FORTRAN and compilers using $-O2$ optimization. The average CPU time for each UHBD simulation was 2150 s, much greater than the time required for the SMOL SSSE calculations.

To compare the results from SSSE and BD calculations for the mAChE system (Tara et al., 1998), the binding rate constants for surfaces 1–6 and ionic strengths (0–670 mM) were plotted in Fig. 8. This comparison shows that most of the SSSE calculations give smaller reaction rates than BD simulations and therefore lead to the disagreement between the methods in the choice of the best reactive surface. There are multiple reasons for this disagreement. Although both the particle-based and continuum methods have the same underlying physics, they employ very different implementations. First and foremost, the particle-based approach uses discrete particle displacements and therefore should only give perfect agreement with the continuum simulations in the limit of infinitesimal step sizes. Second, BD simulations enforce reflective boundary conditions (compare to Eq. 7) only approximately by simply rejecting any steps which take particles into nonreactive surfaces. Given the obvious role of the reflective biomolecular surface in the gorge, it is not surprising that there are significant differences between BD and SSSE results. Finally, we have not implemented any of the error-based adaptive refinement methods available for finite element methods (Axelsson and Barker, 1984; Braess, 1997).

Analysis of SSSE reaction rates and comparison with experiment

The variation of reaction rates with ionic strength is often interpreted via the Debye-Hückel limiting law by Radic et al. (1997),

$$k_{\text{on}} = (k_{\text{on}}^0 - k_{\text{on}}^{\text{H}})10^{-1.18|z_{\text{E}}z_{\text{i}}|\sqrt{I}} + k_{\text{on}}^{\text{H}}, \quad (19)$$

where k_{on} is the observed binding rate constant, k_{on}^{H} is the effective infinite ionic strength limiting rate, k_{on}^0 is the effective 0 ionic strength rate, z_{E} is the effective enzyme

FIGURE 6 Reactive surfaces 1–6 of mAChE (*bottom to top*) from the reactive boundary definition described in the text. The views of these surfaces start from the outside of the protein (*bottom*) and move to the interior (*top*) as the surfaces move inside the active site gorge.

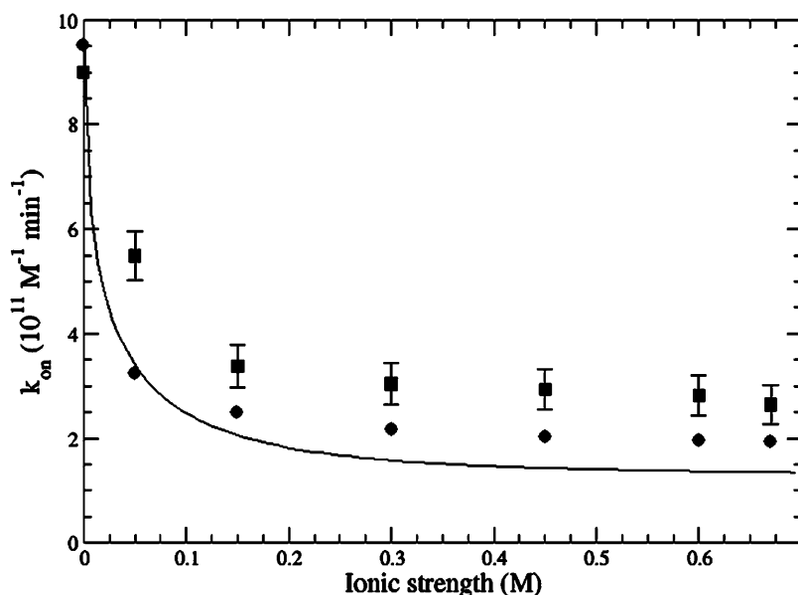


FIGURE 7 Reaction rates of mAChE calculated with the SMOL FE program with reactive surface 1 (●), calculated from Brownian dynamics with reactive surface 1 (■), and from experimental data (Radic et al. 1997) fit to the Debye-Hückel limiting law (solid line).

charge, z_1 is the effective inhibitor charge, and I is the ionic strength. Following the analysis of Radic et al. (1997) of their experimental data, nonlinear regression analysis was used to fit the calculated results at concentrations between 0.050 and 0.450 M and obtain values for k_{on}^0 and z_E . The value of z_1 was held fixed at $+1 e$ and k_{on}^H was set to the value of k_{on} calculated at 0.670 M ionic strength. The results of the rate calculated and the fit parameters are given in Table 2. The data show that the calculated rates fit Eq. 19 very well over the ionic strength range for which the Debye-Hückel limiting law is valid (0.05 M to 0.5 M).

Table 2 shows that the rate constants for reactive surfaces 3–6 do not decrease monotonically with the decreasing reactive patch size. This interesting behavior appears to be influenced by the biomolecular electrostatics; the expected monotonic decrease is observed when the ligand charge is set to 0. A number of residues influence the movement of the ligand within the gorge. Previous work (Tara et al., 1998) showed that residue Asp-74, immediately below reactive surface 3 and above surface 4, plays an important role for the ligand kinetics within the gorge and is a likely candidate for the source of this behavior.

When performing computational diffusion simulations, both particle-based and continuum methods need to choose the appropriate reaction criterion for calculation binding rate constants. This choice is usually made by comparing the simulation results with available experimental data. Fig. 7 indicates that SSSE-derived reaction rates show the best overall agreement with experiment for surface 1. Table 2 also shows that, whereas the calculated results consistently underestimate k_{on}^0 , the best agreement is observed for surface 1. The results obtained at surfaces 3–6 are somewhat less than the experimental values but do show good agreement with one another, indicating that the value of k_{on}^0 is insensitive to the choice of surface definition in this region. Likewise, surfaces 3–6 all give similar values for the effective enzyme charge z_E ; however, unlike k_{on}^0 , the values of z_E from surfaces 3–6 all agree well with experiment. As for k_{on}^0 , reactive surface 1 gives a calculated value, z_E , which matches experimental observations.

The above results show that the SSSE gives very good agreement with experiment when the correct reaction criterion is used. In the case of mAChE, this criterion corresponds to the reactive boundary located at surface 1.

TABLE 2 Rate constants for association of the TFK⁺ inhibitor with mAChE in media of varying ionic strength

Results source	Reactive surface	k_{on}^0 ($10^{11} \text{ M}^{-1} \text{ min}^{-1}$)	k_{on}^H ($10^{11} \text{ M}^{-1} \text{ min}^{-1}$)	$k_{\text{on}}(\text{TFK}_0)$ ($10^{11} \text{ M}^{-1} \text{ min}^{-1}$)	Z_E (e)
Experiment		9.8 ± 0.6	1.3	2.2 ± 0.3	-2.3 ± 0.2
SMOL results	1	6.79 ± 0.45	1.93	1.42	-2.07 ± 0.07
	2	3.82 ± 0.39	0.15	0.38	-1.16 ± 0.08
	3	2.87 ± 0.55	0.02	0.025	-1.89 ± 0.15
	4	4.79 ± 0.98	0.026	0.017	-1.98 ± 0.16
	5	5.01 ± 1.03	0.025	0.0077	-2.00 ± 0.16
	6	5.59 ± 1.17	0.028	0.0061	-2.00 ± 0.17

The various parameters were obtained by fitting to Eq. 19 using fixed values k_{on}^H fixed at the measured/calculated k_{on} at 0.67 M ionic strength and Z_1 fixed at $+1 e$. The $k_{\text{on}}(\text{TFK}_0)$ column denotes direct calculations for the ligand with zero charge. Experimental results are from Radic et al. (1997).

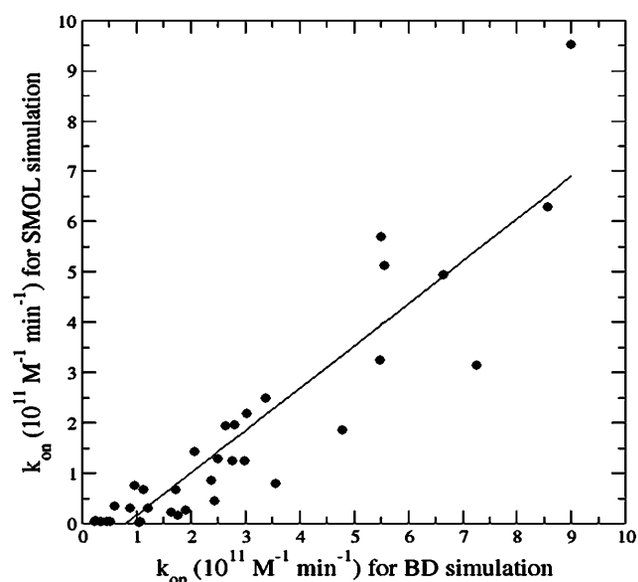


FIGURE 8 Correlation plots for calculations of reaction rate using the SMOL program and using BD with spherical surface reactive boundary definitions. For each method, calculations are made for six reactive surfaces and ionic strength ranges 0.00, 0.05, 0.10, 0.15, 0.3, 0.45, 0.6, and 0.67 M. For ● data points, BD results are plotted on the x axis, SMOL results are plotted on the y axis. The solid line is the linear fit of the data, and its slope is 0.84 with a Pearson correlation coefficient of 0.91.

The BD results by repeating the simulation of Tara et al. (1998) also used spherical reaction criteria similar to those used in the present study. However, the reaction rates results suggest that surface 3 provides the best agreement with experiment. The differences between BD and SSSE results were discussed above (see also Fig. 8). It is not surprising that the two methods give different rates at the same surfaces; it is encouraging that both methods offer reaction conditions which give good agreement with the experimental results.

CONCLUSIONS

In this study, we describe new continuum-based methods for studying diffusion in biomolecular systems. Specifically, we present the SMOL software package, a finite element-based software package for solving the SSSE to calculate ligand binding rate constants for large biomolecules. Additionally, we describe new adaptive meshing methods developed to discretize biomolecular systems into finite element meshes which respect the geometry of the biomolecule. Although not presented in this study, it is important to note that the new meshing methods could be useful in a variety of biological simulations including computational studies of biomolecular electrophoresis (Allison, 2001), elasticity (Zhang et al., 2003a), and electrostatics (Baker et al., 2000; Cortis and Friesner, 1997a,b; Holst et al., 2000). The SMOL software was validated using a spherical analytical test case. Rates for inhibitor binding to mAChE were calculated at various ionic strengths with several different reaction criteria. The cal-

culated rates were compared with experimental data (Radic et al., 1997) and show very good agreement with experiment while requiring substantially less computational effort than existing particle-based Brownian dynamics methods.

Since the inception of this work, new Brownian dynamics methods have been developed which dramatically improve the speed and convergence of these particle-based simulations (Zou and Skeel, 2003). However, one of the key ingredients to these new methods is the construction of an a priori approximate probability distribution to bias and accelerate the BD calculations. One interesting future direction of the present research would be the use of these continuum methods to develop approximate biasing functions for the new biased BD methodology of Zou and Skeel.

This initial research lays the groundwork for several new directions. Of particular interest is the integration of molecular-scale information into simulations of cellular-scale systems such as the neuromuscular junction (Smart and McCammon, 1998; Tai et al., 2003). Additionally, this new finite element framework should facilitate the incorporation of other continuum mechanics phenomena into biomolecular simulations. The ultimate goal of this work is to develop scalable methods and theories that will allow researchers to begin to study biological macromolecules in a cellular context.

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